Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
S1	1	10/521782	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 10:22
S2	16979	epoxidation or oxidation and "tube reactor"	US-PĞPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	·OR	ON	2007/08/29 18:47
<b>S3</b>	3004	(epoxidation or oxidation) and "tube reactor"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/29 18:48
<b>S4</b>	171	epoxidation and "tube reactor"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/29 18:49
S5	106	S4 and (hydroperoxide or "hydrogen peroxide")	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/29 18:51
S6	502	S3 and (hydroperoxide or "hydrogen peroxide")	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/29 19:12
	28	S6 and "feed point"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 10:01

S8	3013	(epoxidation or oxidation) and "tube reactor"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 10:01
S9	. 503	S8 and (hydroperoxide or "hydrogen peroxide")	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 10:01
S10	<b>44</b>	S9 and adiabatic	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 10:02
S11	, 616	reactor and "tube reactor" and "fixed-bed"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 13:02
S12	39	S11 and "feed points"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 10:37
S13	. 122	S11 and vertical	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 11:22
S14		S8 and ("two feed points" or "dual feed")	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 11:30

S15	12	S8 and ("after-reactor")	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR .	ON	2007/08/30 11:33
S16	5	"wo 01/10855"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR .	ON	2007/08/30 11:56
S17	6	"wo 01/72729"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR .	ON	2007/08/30 13:00
S18		"wo 02/22259"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 13:00
S19	371	"tube reactor" and ("feed points" or "feed inlets" or "feed ports")	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 13:04
S20	9	"4582613"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:18
S21	5	"6410786"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:21

S22	7	"6563000"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:22
S23	7	09/387119	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:23
S24	1,	10/387119	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:23
S25	2	"6949675"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:55
S26	425	549/524 or 549/529	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:56
S27	21	S26 and "tube reactor"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/08/30 14:57

Ref #	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L2	438	549/531	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:08
L3	0	I2 and reactor and "inlet points" and feed	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:09
L4	2	I2 and oxirane and hydroperoxide and catalyst and adiabatic and ((tube or "fixed-bed") near reactor)	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:09
L5	0	I2 and reactor and "multiple feed points"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:10
L6	0	I2 and reactor and "inlet points" and feed	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:10
L7	28	I2 and "tube reactor"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR ·	ON	2007/10/01 17:10
L8 .	8	I2 and (oxirane or "propylene oxide") and (hydroperoxide or "hydrogen peroxide" or hydrogenperoxide) and catalyst and adiabatic and ((tube or "fixed-bed") near reactor) and zeolite and solvent	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:21

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S1	1	10/521782	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:08
S2	13	oxirane and hydroperoxide and catalyst and adiabatic and tube and reactor	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 13:58
S3	. 10	oxirane and hydroperoxide and catalyst and adiabatic and ((tube or "fixed-bed") near reactor)	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:09
S4	31	(oxirane or "propylene oxide") and (hydroperoxide or "hydrogen peroxide" or hydrogenperoxide) and catalyst and adiabatic and ((tube or "fixed-bed") near reactor) and zeolite and solvent	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:21
S5	2	10/220503	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 14:51
S6	63	reactor and "multiple feed points"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:10
S7	1	S6 and epoxidation	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 14:54

			r		·	
S8	410	reactor and "inlet points" and feed	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:10
S9	11	S8 and epoxidation	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 14:54
S10	3138	reactor and epoxidation	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 15:11
S11	1053	S10 and hydroperoxide	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON .	2007/10/01 15:11
S12	1000	S11 and catalyst	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 15:12
S13	783	S12 and (propene or propylene)	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 15:14
S14	73	S13 and "tube reactor"	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT; IBM_TDB	OR	ON	2007/10/01 17:10

IBM_TDB	S15	13	S14 and adiabatic	US-PGPUB; USPAT; USOCR; FPRS; EPO; JPO; DERWENT;	OR	ON	2007/10/01 15:16
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10/1/2007 5:22:25 PM C:\Documents and Settings\Dgallis\My Documents\EAST\Workspaces\10-521782b.wsp

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L1

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FILE 'CAPLUS' ENTERED AT 17:36:27 ON 01 OCT 2007

O S (OXIRANE OR "PROPYLENE OXIDE") AND (HYDROPEROXIDE OR "HYDROGE

L2	0	s	(OXIRANE OR "PROPYLENE OXIDE") AND (	HYDROPEROXIDE OR "H	HYDROGE
L3	1	s	(OXIRANE OR "PROPYLENE OXIDE") AND (F	HYDROPEROXIDE OR "H	HYDROGE
L4	1414	s	(OXIRANE OR "PROPYLENE OXIDE") AND (F	HYDROPEROXIDE OR "H	HYDROGE
L5	4	S	L4 AND CATALYST AND ADIABATIC		
L6	103	S	L4 AND CATALYST AND REACTOR		

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L7 20 S L6 AND (TUBE OR "FIXED-BED")
L8 1 S L7 AND SOLVENT AND ZEOLITE

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=> d 17 1-20 abs ibib

ANSWER 1 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN Gas-propylene was epoxidized with dilute methanol solns. of hydrogen peroxide in a fixed-bed reactor using self-made titanium-containing mol. sieve of ZSM-5 type (Model

using self-made titanium-containing moi. sleve of 2010-2019 to 2010-2019 as catalyst. The technol. conditions such as the liquid space-velocity, mole ratio of propylene to hydrogen peroxide, temperature and contents of hydrogen peroxide in methanol solns. which would determine the epoxidn. rate, were studied systematically. The exptl. results show that when the reaction pressure is 0.7 MPa, reaction temperature is 50 °C, mass content of hydrogen peroxide in methanol solns. is about 1.5%, liquid mass space velocity is 48 h-1, the conversion of hydrogen peroxide and selectivity of propylene oxide are more than 901 and 85%, resp.

ISSION NUMBER: 2007:1065469 CAPLUS

Electric Epoxidation of propylene and hydrogen peroxide in a lower pressure continuous fixed-bed reactor

CRC(S): 2010-2019 Shen, Ben-xian; Xiao, Wei-guo; Zhang,

ACCESSION NUMBER:

AUTHOR (S):

fixed-bed reactor
Zhao, Jl-gang; Shen, Ben-Xian; Xiao, Wei-guo; Zhang,
Ce; Wang, Lei
School of Chemical Engineering, East China University
of Science and Technology, Shanghai, 200237, Peop.
Rep. China
Huadong Ligong Daxue Xuebao, Ziran Kexueban (2007),
33(11), 23-27
CODEN: HIXEEV; ISSN: 1006-3080
Huadong Ligong Daxue Xuebao Bianjibu
Journal CORPORATE SOURCE:

SOURCE:

PUBLISHER: DOCUMENT TYPE:

LANGUAGE: Chinese

CN 1546479 PRIORITY APPLN. INFO.:

A

L7 ANSWER 3 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
AB Hydrogen and oxygen react in a plasma generating apparatus of medium barrier discharging at ambient temperature and atmospheric to give hydrogen peroxide, which is gathered with methanol and used directly in the epoxidn. of propylene. The plasma generation apparatus is a coaxial glass structure and the circulation cooling water in the annular space of the sleeve also acts as an earth electrode. The epoxidn. of propylene is carried out in batch or fixed bed reactor.

ACCESSION NUMBER: 2005:561306 CAPLUS

DOCUMENT NUMBER: 143:13916

Wethod and device for epoxidation of propylene using plasma of hydrogen and oxygen

GUO, Hongchen

PATENT ASSIGNEE (S): Dalian University of Technology, Peop. Rep. China Faming Zhuanii Shenqing Gongkai Shuomingshu, No pp. given

DOCUMENT TYPE: Patent DOCUMENT TYPE: LANGUAGE: . FAMILY ACC. NUM. COUNT: PATENT INFORMATION: KIND DATE APPLICATION NO. DATE

20041117

CN 2003-10105211 CN 2003-10105211

20031128

ANSWER 2 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
A process for the epoxidn. of olefins with hydrogen
peroxide is claimed. This process serves to increase product
yield by minimizing loss of olefin conversion rate and inhibit
fuction of
byproducts while using a catalyst formed by a conventional
method. Said method serves to improve the inactivation degree of the
catalyst in a continuous reaction. The epoxidn of alkenes is
performed by a reaction with hydrogen peroxide in the
presence of titanium silicalite catalyst (formed by using
alumina as a binder or a supporter) with olefin in a fixed
bed reactor using an alc.-nitrile mixed solvent. The
titanium silicalite catalyst formed is a crystalline titanium-containing
zeolite having the NFI or MEL structure represented by the formula
xTiO2(1-x)SiO2 (x = 0.01-0.05). The titanium silicalite catalyst
formed is a mixture of titanium silicalite and 5-95 weight of the alumina
binder or support. The nitrile is acetonitrile, propanenitrile or
pentanenitrile and the alc. is methanol, ethanol, isopropanol,
tett-butanol, cyclohexanol. The amount of hydrogen
peroxide is 1-90 weighth: The olefin is a C2-10-aliphatic olefin. The
epoxidn. is performed at 0-100° under 1-100 atm pressure and the
mass-space velocity is 0.05-1 h-1. More narrow definitions are
indicated. however, specific chemical structures and/or addnl. information are not provided here.

ACCESSION NUMBER: 2006:1141411 CAPLUS

DOCUMENT NUMBER: 146:295762 146:295762
Process for epoxidation of olefin with
hydrogen peroxide and titanium
silicalite catalyst
Kwak, Byong Sung: Oh, Seung Hoon: Lee, Jae Ho; Choi,
Hyun Chul: Kim, Tae Jin; Lee, Jong Hyung: Choo, Dae TITLE: INVENTOR(S): Hyun Sk Corporation, S. Korea Repub. Korean Kongkae Taeho Kongbo, No pp. given CODEN: KRXXA7 PATENT ASSIGNEE(S): SOURCE: DOCUMENT TYPE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE KR 2006055409 PRIORITY APPLN. INFO.: А 20060523 KR 2005-110355 20051117

ANSWER 4 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
Epoxidn. of propylene to propylene oxide over Ag/TS-1
in the presence of oxygen and hydrogen was carried out in a fixed
-bed, quartz flow reactor. The effects of Ag loading,
support, Si/Ti mol. ratio of TS-1 and calcination temperature of Ag/TS-1

ne propylene epoxidn. were investigated. The results show that Ag loading, support and the calcination temperature of Ag/TS-1 as well as the Si/Ti

ratio of TS-1 have a great effect on the catalytic properties. The optimum Ag loading, calcination temperature of Ag/TS-1 and Si/Ti mol. ratio of

ratio of

TS-1 are 2 weights, 450°C and 64, resp. Over 2.0 weights Ag/TS-1
(nSis/nTi=64) catalyst, at a space velocity of 3000h-1, 0.92's
propylene conversion with 91.2's selectivity to propylene
oxide is obtained at 150°C. The descrivation of Ag/TS-1
catalyst is not due to the changes of active species, but the coke
of the catalyst, which can be easily regenerated by calcination
at 450°C in air.

ACCESSION NUMBER:

2005:418395 CAPLUS
DOCUMENT NUMBER:
144:150677
TITLE:
Propylene envidation over Ag/TS-1 catalysts

AUTHOR(S): CORPORATE SOURCE:

144:150617
Propylene epoxidation over Ag/TS-1 catalysts
Wang, R.; Hao, J.; Guo, X.; Wang, X.; Liu, X.
State Key Laboratory of Fine Chemicals, Dalian
University of Technology, Dalian, 116012, Peop. Rep.
China
Studies in Surface Science and Catalysis (2004),
154C(Recent Advances in the Science and Technology of
Zeolites and Related Materials), 2632-2638
CODEN: SSCTDM; ISSN: 0167-2991
Elsevier B.V.
Journal
English
13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR

PUBLISHER: DOCUMENT TYPE: LANGUAGE: REFERENCE COUNT: THIS

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SOURCE:

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ANSWER 5 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
Synthesis of propylene oxide through gas phase
epoxidn. of propylene using mol. oxygen as an oxidant over lab-prepared
Ag/TS-1 catalyst and the lab fixed-bed
reaction unit with quartz reactor was studied. The preparation
conditions of Ag/TS-1 catalyst such as, Ag loading, type of
support, the Si/Ti mol ratio of TS-1, calcination conditions, etc. were
investigated. The effects of reaction conditions including reaction
temperature, gas velocity, volume ratio of C3H6/O2 and C3H6/H2 on
lyst
                                                                                                                                                                                                                                                                                                                                                                                    ANSWER 6 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN The deactivation and regeneration of the lamina titanium silicalite
                                                                                                                                                                                                                                                                                                                                                                                    atalyst for the epoxidn. of propylene with dilute H2O2 was investigated in a fixed-bed reactor. In the scale-up experiment, the dosage of the lamina TS-1 catalyst is 2.5 kg, after 1000 h reaction the catalyst still exhibits good performance and further increases the reaction time, the conversion of H2O2 begins to decrease. TG and BET analyses of the deactivated catalysts show that the main species occluded within the zeolite pore are propylene oxide oligomers, and these species occupying the active Ti site and blocking the pores of the lamina TS-1
catalyst

performance were also investigated. The optimum Ag loading w(Ag) was of 21 (mass percent) and n(Si)/n(Ti) was of 64 (support), and the optimum calcination temperature was 450°C in air for Ag/Ts-1 catalyst. The reaction when carried out at 150°C by passing the reactant mixture of C3H6, O2, H2 and N2 through the reactor at a volume ratio of 1:2:3:12 with a space velocity of 4000 h-1 is optimum. After 70 min reaction, propylene conversion (x(C3H6)) and selectivity to propylene oxide (SPO) were 1.374 and 93.513, resp.

ACCESSION NUMBER: 2004.752970 CAPLUS

DOCUMENT NUMBER: 142:156346

TITLE: Propylene oxidation to propylene
                                                                                                                                                                                                                                                                                                                                                                                      the main reason for the deactivation of catalyst. The deactivated catalyst can be regenerated by different regeneration methods. The activity of deactivated catalysts regenerated by dilute H202 or heat treatment by using air or nitrogen as
                                                                                                                                                                                                                                                                                                                                                                 calcination media can be fully recovered, but a decline in propylene oxide (PO) selectivity of the regenerated catalyst has been observed during the first hours of reaction. However, water vapor treatment of the deactivated catalyst can improve the PO selectivity with the same activity as that of the fresh lamina TS-1 catalyst.

ACCESSION NUMBER: 141:351728
                                                                                                 142:156346

Propylene oxidation to propylene oxide over Ag/TS-1 catalyst in the gas phase wang, Rui-pu: Rao, Jing-quan; Guo, Xin-wen; Wang, Xiang-sheng; Liu, Xue-wu State Key Laboratory of Fine Chemicals, Dalian University of Technology, Dalian, 116012, Peop. Rep. China Shiyou Xuebao, Shiyou Jiagong (2004), 20(4), 44-50 CODEN: SXSHEY; ISSN: 1001-8719
Shiyou Xuebao, Shiyou Jiagong Bianjibu Journal Chinese
  TITLE:
  AUTHOR (S):
  CORPORATE SOURCE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Regeneration of Lamina TS-1 Catalyst in the Epoxidation of Propylene with Hydrogen
                                                                                                                                                                                                                                                                                                                                                                   TITLE:
  SOURCE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Feroxide
Liu, Xuewu; Wang, Xiangsheng; Guo, Xinwen; Li, Gang;
Yan, Haisheng
State Key Laboratory of Fine Chemicals, Dalian
University of Technology, Dalian, 116012, Peop. Rep.
                                                                                                                                                                                                                                                                                                                                                                  AUTHOR (S):
   PUBLISHER:
DOCUMENT TYPE:
LANGUAGE:
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                     China
Catalysis Letters (2004), 97(3-4), 223-229
CODEN: CALEER; ISSN: 1011-372X
Kluwer Academic/Plenum Publishers
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DOCUMENT TYPE:
LANGUAGE:
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CASREACT 141:351728
28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR
                                                                                                                                                                                                                                                                                                                                                                    OTHER SOURCE(S):
                                                                                                                                                                                                                                                                                                                                                                    REFERENCE COUNT:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           RECORD. ALL CITATIONS AVAILABLE IN THE RE
                                                                                                                                                                                                                                                                                                                                                                   FORMAT
ANSWER 7 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN

AB The TS-1 powder, which is synthesized using tetrapropylammonium bromide
(TPABr) as template, was molded into catalyst by extruding. The
prepared catalysts containing 301, 501, 601 or 801 TS-1 were used to
catalyze the propylene epoxidn. to produce PO in a fixed
bed reactor. The results show that TS-1 is the active
site of the catalyst. The catalyst with 801 TS-1 has
the highest activity and mach. Intensity. The selectivity of PO
decreases
and the utilization of hydrogen peroxide increases as
TS-1 content increases in the catalyst. Both the selectivity of
PO and the utilization of hydrogen peroxide increase
as the WHSV of propylene increases from 0.30 h-1 to 0.70 h-1. The
catalyst containing 801 TS-1 exhibits 97.81 hydrogen
peroxide conversion, 86.41 PO selectivity and 91.81
hydrogen peroxide utilization when the WHSV of propylene
increases (COM 10.0) held (COM 10.0) held (COM 10.0) held
COLUMENT NUMBER: 2004:365388 CAPLUS
DOCUMENT NUMBER: 141:157499

**TITLE:** Properties of titanium silicalite catalyst
with different TS-1 contents
                                                                                                                                                                                                                                                                                                                                                                                    ANSWER 8 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
The present invention relates to a process for the epoxidn. of ≥1
organic compound with an oxygen-delivering substance, for example a
hydroperoxide, in the presence of ≥1 catalyst
containing a metal-organic framework material comprising pores and a
                                                                                                                                                                                                                                                                                                                                                                                      and \geq 1 bidentate organic compound, said bidentate organic compound being coordinately bound to the metal ion. Thus, a 66:24:10 volume ratio of
                                                                                                                                                                                                                                                                                                                                                                    O2,

He, and propylene was streamed through a tube reactor containing AgMO3-treated MOF-5 at 220° to give propylene oxide with a turnover of 3.3% and selectivity of 10.3% ofter 15 h. ACCESSION NUMBER: 2003:747904 CAPLUS
                                                                                                                                                                                                                                                                                                                                                                    DOCUMENT NUMBER:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     139:278233
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    139:278233
Process for epoxidation of organic compounds with oxygen or oxygen-delivering compounds using catalysts containing metal-organic framework
                                                                                                   2004:369388 CAPLUS
141:157499
Properties of titanium silicalite catalyst
with different TS-1 contents
Song, Gang; Li, Gang; Guo, Xin-wen; Wang,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    catalysts containing metal-organic framework (MOF) materials
Mueller, Ulrich: Lobree, Lisa; Hesse, Michael; Yaghi,
Omar M.: Eddaoudi, Mohamed
BASF Aktiengesellschaft, Germany; The Regents of the
University of Michigan
U.S., 13 pp.
CODEN: USXXAM
                                                                                                                                                                                                                                                                                                                                                                    INVENTOR(S):
                                                                                                                                                                                                                                                                                                                                                                    PATENT ASSIGNEE (S):
                                                                                                                                                                                                                                                                                                                                                                    SOURCE:
                                                                                                     Chen, Yong-ying
Department of Catalytical Chemistry and Engineering,
Dalian University of Technology, Dalian, 116012,
  CORPORATE SOURCE:
                                                                                                                                                                                                                                                                                                                                                                    DOCUMENT TYPE:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     English
                                                                                                                                                                                                                                                                                                                                                                    FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
   Peop.
                                                                                                    Rep. China
Ranliao Huaxue Xuebao (2004), 32(1), 113-116
CODEN: RHXUD8; ISSN: 0253-2409
Kexue Chubanshe
Journal
Chinese
   SOURCE:
                                                                                                                                                                                                                                                                                                                                                                                      PATENT NO.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                     KIND
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DATE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              APPLICATION NO.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DATE
    PUBLISHER:
     DOCUMENT TYPE:
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20030923
20031211
                                                             US 2002-157494
WO 2003-EP5547
                                                                                             20020530
      US 6624318
                                   B1
A1
       WO 2003101975
      MO 2003101975 A1 2003223 W: US
RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
IT, LU, MC, NL, PT, RO, SE, SI, SK, TR
EP 1513823 A1 20050316 EP 2003-730125 20030527
EP 1513823 B1 20070808
      AT 369350
PRIORITY APPLN. INFO.:
                                                             WO 2003-EP5547
                                                                                        W 20030527
```

OTHER SOURCE(S): REFERENCE COUNT:

CASREACT 139:278233
5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

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L7 ANSWER 9 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) RECORD. ALL CITATIONS AVAILABLE IN THE RE
   L7 ANSWER 9 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
AB A process for the production of an epoxide (e.g., methyloxirane) by
reaction
AB A process for the production of an epoxide (e.g., methyloxirane) by reaction

of an olefin (e.g., propylene) with a hydroperoxide (e.g., benzene hydroperoxide) in the presence of a solid epoxidn.

catalyst is described where the olefin and hydroperoxide reactants are first reacted in it least two fixed-bed reactors packed with fresh catalyst, olefin being fed aerially to the reactors and hydroperoxide reactant being fed in parallel to the reactors and then the reactor effluent from the last of the reactors is passed to as least one further fixed-bed reactor packed with epoxidn. catalyst which has been at least partially deactivated. A process flow diagram is presented.

ACCESSION NUMBER: 2003:485738 CAPLUS

DOCUMENT NUMBER: 139:53455

TITLE: Epoxidation system with fixed-bed reactors for the manufacture of epoxides from alkenes and hydroperoxides

Leyshon, David W.; Jubin, John C., Jr.; Wolff, Richard
                                                                                                                                                                                                                                                                                                                                                                                    FORMAT
   INVENTOR(S):
Richard
                                                                                                        U.S., 5 pp.
CODEN: USXXAM
   PATENT ASSIGNEE(S):
   DOCUMENT TYPE:
                                                                                                         Patent
English
              IGUAGE:
   FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                 EP 1551819

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
BR 2003015277

A 20050830

BR 2003-15277

A 20050830

BR 2003-15277

A 20050830

BR 2003-15277

A 20050830

BR 2003-15277

C 20030827

AT 3188052

T 20060315

AT 2003-824268

20030827

AT 2006513155

T 20060320

JP 2006-345255

20030827

BES 2254987

T3 20060616

ES 2254987

BY 2006-345255

20030827

BY 2006-345255

BY 2006-345256

BY 2007-345268

BY 2007-34
   BR 20030132/7
CN 1688562
AT 318802
JP 2006513155
ES 2254987
PRIORITY APPLN. INFO.:
                                                                                                                                                                                       WO 2003-US27158
                                                                                                                                                                                                                                                                        W . 20030827
                                                                                                         MARPAT 139:53455
   OTHER SOURCE(S):
                                                                                                                                  THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
   REFERENCE COUNT:
                     ANSWER 10 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
Titanium silicalite (TS-1) was successfully synthesized by using Pr4NBr
(TPABr) and BuNH2 as the template and the base, resp. XRD, IR, SEM, 13C
CP/MAS NMW, UV-visible, and UV-Raman techniques were used to characterize
the synthesized products. The results show that TS-1 has MFI structure
with high crystallinity and large crystal size, and with two kinds of
titanium species. Using epoxidn. of propylene as the probe, the
                       lytic
properties of TS-1 in the present work were investigated. The crystal
size of TS-1 greatly affects the initial reaction rate of
propylene oxide (PO) formation. The effects of reaction
conditions on PO reveal that the PO formation is greatly influenced by
                         solvent and the solution pH value. Methanol is the most preferable
     solvent,
and a portion of water (<30%) in solution does not have much impact on
                       reaction. Moreover, the addition of base used to neutralize the residual acidity of the catalyst favor the reaction, the catalyst shows high activity and selectivity upon the adequate amount of base and
  catalyst can be used repeatedly without regeneration. In the
fixed-bed reactor, using ammonia water as the
neutralizer, after 200 h of reaction, both the conversion and the
utilization of H2O2 is about 951, the selectivity of PO is about 901.
ACCESSION NUMBER: 2002-C403022 CAPIUS
DOCUMENT NUMBER: 137:232944
TITLE: Synthesis of titanium silicalite (TS-1) from the
                                                                                                          system and its catalytic properties for epoxidation
                                                                                                         propylene
Wang, Xiang-sheng; Guo, Xin-wen; Li, Gang
State Key Laboratory of Fine Chemicals, Dalian
University of Technology, Dalian, 116012, Peop. Rep.
     AUTHOR(S):
CORPORATE SOURCE:
                                                                                                          China
Catalysis Today (2002), 74(1-2), 65-75
CODEN: CATTEA; ISSN: 0920-5861
Elsevier Science B.V.
    SOURCE:
                                                                                                                                                                                                                                                                                                                                                                                                          PATENT NO.
    PUBLISHER:
DOCUMENT TYPE:
      LANGUAGE:
REFERENCE COUNT:
                                                                                                          English
27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR
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RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

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ANSWER 11 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
Propylene oxide is manufactured in a three-stage process
from propylene, oxygen, and hydrogen. The first reaction step is the
oxidation of isopropanol/water with mol. oxygen in a
tion-distillation column
(approx. 500 psi and 350° F.), to produce hydrogen
peroxide and acetone. The column is configured with an upper high
liquid holdup reaction zone and a lower short residence time stripping
                                 Inert gas circulating through the column effects separation of the hydrogen peroxide as part of the bottoms fraction and acetone as part of the distillate fraction. The liquid part of the distillate fraction comprising acetone, isopropanol and water is then reacted with hydrogen (second reaction step) under reactive-distillation conditions to convert the contained acetone back to isopropanol for subsequent recycle to the first reaction step. The third reaction step
                                 the epoxidn. of propylene (in stoichiometric excess) with the hydrogen peroxide solution, typically in the presence of a titanium silicalite catalyst. The reaction is performed in a series of fixed bed adiabatic reactors with
intercooling. Product separation is by conventional distiliation
Unceated
propylene is recycled to the epoxidn. step and water/isopropanol to the
first reaction step.

ACCESSION NUMBER: 2002:23864 CAPLUS
DOCUMENT NUMBER: 136:70254
Three-stage process for manufacturing of
propylene oxide
Gelbein, Abraham P.
Chemical Research & Licensing Company, USA
SOURCE: U.S., 9 pp.
CODEN: USXXAN
DOCUMENT TYPE: Patent
LANGUAGE: Patent
LANGUAGE: Patent
English
FAMILY ACC. NUM. COUNT:
DATENT INFORMATION:
                                                                                                                                                                                                                                                                                APPLICATION NO.
                                                                                                                                                                                                                                                                                                                                                                                                                              DATE
   W1 6337412 B1 20020108 US 2001-841427 20010424
W2 2002085876 A1 20021031 W0 2001-US49838 20011228
W1 AE, AG, AL, AM, AT, AL, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CC, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GK, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, M, MW, MV, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW
RN: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CN, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
AU 2002245164 A1 20021105 AU 20002-245164 20011228
PRIORITY APPLN: INFO::
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THERE ARE 17 CITED REFERENCES AVAILABLE FOR RECORD. ALL CITATIONS AVAILABLE IN THE RE

REFERENCE COUNT:

FORMAT

ANSWER 12 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN

The title process comprises passing a feed containing an organic hydroperoxide, e.g., ethylbenzene hydroperoxide, and alkene, e.g., propene, through a bank of ≥2 serially connected reactors all containing a bed of heterogeneous epoxidn. catalyst particles, e.g., com. titanis/alumina, and operated in a cyclic mode. The lat reactor of the cyclically operated bank is put in a position further down this bank, when the activity of the epoxidn. catalyst contained in the reactor decreased to an undesirably low level. In this position the catalyst with decreased activity is contacted with the effluent from the reactor in the preceding position at a temperature which is ≥5\* higher than the final temperature at which the catalyst was in use in the lst position of the bank and for sufficient time to restore its activity to the desired level. The bank of cyclically operated epoxidn. reactors may be followed by ≥1 addnl. fixed-bed epoxidn. reactors which are not operated cyclically.

ACCESSION NUMBER: 2001:137204 CAPLUS
DOCUMENT NUMBER: 134:194867
TITLE: Process for preparing alkylene exides

TITLE: INVENTOR (S):

Process for preparing alkylene oxides Derks, Willen; Dirkzwager, Hendrik; Van der Veen, Alexander Jan; Wermeling, Rutger Johannes Franciscus Shell Internationale Research Maatschappij BV, Neth. PATENT ASSIGNEE (S):

PCT Int. Appl., 25 pp. CODEN: PIXXD2 Patent SOURCE:

DOCUMENT TYPE:

LANGUAGE: Facent English FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

PAT	PATENT NO.									APPLICATION NO.							DATE		
WO	2001						2001	0222			2000-				. 2	0000	816		
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	ВВ	, BG,	BR,	BY,	BZ,	CA,	CH,	CN,		
		CR,	CU,	C2,	DE.	DK,	DM,	EE,	ES,	FI	, GB,	GD,	GE,	GH,	GM,	HR,	HU,		
		ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR	, K2,	LC,	LK,	LR,	LS,	LT,	LU,		
		LV,	MA.	MD,	MG,	MK.	MN,	MW,	MX,	MZ	, NO.	NZ,	PL.	PT,	RO,	RU,	SD,		
		SE.	SG,	SI.	SK,	SL.	TJ,	TM,	TR,	TT	, TZ,	UA,	UG,	US,	UZ,	VN,	YU,		
		ZA,	ZW																
	RW:	GH,	GM,	KE,	LS,	MW.	MZ,	SD,	SL,	SZ	, TZ,	UG,	ZW.	AT.	BE,	CH,	CY,		
											, LU,								
											, NE,								
us	6365				B1						2000-				2	0000	60B		
BR	2000	0133	58		A		2002	0430		BR	2000-	1335	В		2	0000	816		
EP	1204	653			A1		2002	0515		EΡ	2000-	9546	38		2	0000	816		
EP	1204	653			B1		2003	0129											
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	HC,	PT.	IΕ,		
								CY.											
JP	2003	5073	74		т		2003	0225		JP	2001-	5175	15		2	0000	816		
AU	7576	12			B2		2003	0227		ΑU	2000-	6702	4		2	0000	816		
ES	2186	655			Т3		2003	0516		ES	2000- 2000-	9546	38		2	0000	816		
CN	1135	225			В		2004	0121		CN	2000-	8116	61		2	0000	816		
RU	2241	706			C2		2004	1210		RU	2002-	1068	23		2	0000	816		
PRIORITY	APP	LN.	INFO								1999-				A 1	9990	818		
										WO	2000-	EP80	52	1	W 2	0000	816		

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

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(Continued)
L7 ANSWER 12 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
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L7 ANSWER 13 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN

AB Propylene epoxidn. over a Pd-Pt/TS-1 [Ti silicalite] catalyst with in situ formed hydrogen peroxide was carried out in a fixed bed reactor under high pressure.

The continuous operation allowed the study of catalyst deactivation and changes in product distribution with time-onstream. The initial propylene oxide selectivity was high, 991 at 3.51 conversion, but the catalyst deactivated rapidly with time-onstream and successively the formation of Me formate became the prevalent reaction. Using carbon dioxide, instead of nitrogen, had a beneficial effect on the formation of propylene oxide, and even higher yields were obtained when increasing the pressure from 50 to 120 bar (supercrit fluid phase). Thermal anal. (TA-MS and TA-FTIR) indicated that catalyst regeneration requires oxidation at elevated temperature; washing with an organic solvent is less efficient. The serious

catalyst deactivation and the striking shift in the selectivity pattern of the catalyst is traced to competing alc. oxidation on platinum metal.

ACCESSION NUMBER: 2001:92193 CAPLUS

DOCUMENT NUMBER: 134:312764

TITLE: Continuous epoxidation of propylene with oxygen and hydrogen on a Pd-Pt/TS-1 catalyst

AUTHOR(S): Jenzer, Gregor; Mallat, Tamas; Maciejewski, Marek;

2001:92193 CAPLUS
134:312764
Continuous epoxidation of propylene with oxygen and hydrogen on a Pd-Pt/TS-1 catalyst
Jenzer, Gregor; Mallat, Tamas; Maciejewski, Marek;
Eigenmann, Florian; Baiker, Alfons
Laboratory of Technical Chemistry, Swiss Federal
Institute of Technology, ETH-Zentrum, Zurich, AUTHOR (S): CORPORATE SOURCE:

CH-8092,

SOURCE:

Switz.
Applied Catalysis, A: General (2001), 208(1,2), 125-133
CODEN: ACAGE4; ISSN: 0926-860X
Elsevier Science B.V.
Journal
English
32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR PUBLISHER: DOCUMENT TYPE: LANGUAGE: REFERENCE COUNT: THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

L7 ANSMER 14 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN

AB A closed system comprises a shell and tube reactor and
a shell and tube cooler placed side-by-side, a reversed Utube connecting the top of the reactor and cooler, a Utube connecting the bottom of the reactor and cooler, a
gas-liquid separator on the top of the reactor and connecting to
the reversed U-tube. Finely granular T1-S1 mol. sieves (TS 1)
were used as catalysts. The conversion of H202 and the
selectivity (vs. H202) of propylene oxide were >90%.

ACCESSION NUMBER: 2001:55341 CAPLUS
DOCUMENT NUMBER: 134:86625

Appearatus and processes for epoxidizing propy

DOCUMENT NUMBER: TITLE: Apparatus and processes for epoxidizing propylene with

INVENTOR (S):

hydrogen peroxide solutions Xu, Xi'en; Chen, Xiaohui; Li, Shijun China Petrochemical Group Corp., Peop. Rep. China Faming Zhuanli Shenqing Gongkai Shuomingshu, 6 pp. CODEN: CNXEV PATENT ASSIGNEE (S): SOURCE:

DOCUMENT TYPE:

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

APPLICATION NO. PATENT NO. KIND DATE DATE CN 1256273 PRIORITY APPLN. INFO.: A 20000614 CN 1998-123136 CN 1998-123136 19981209

L7 ANSWER 16 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
AB An olefin epoxidn. process is operated using a plurality of
reactor vessels, each containing a fixed bed of a
heterogeneous catalyst such as titania-on-silica. The
reactor vessels are connected in series whereby a feedstream
comprised of olefin and an active oxygen species is passed through the
series of reactor vessels in contact with the heterogeneous
catalyst to accomplish conversion of the olefin to the
corresponding epoxide. As the activity of the catalyst in an
individual reactor vessel falls to an undesirably low level, the
reactor vessel is taken out of service and a replacement
reactor vessel ontaining fresh or regenerated catalyst
introduced. The replacement reactor vessel may, in alternative
embodiments of the process, be the first or the last reactor
vessel in the series. For example, the feedstream may first be contacted
with either the most active or the least active charge of catalyst
within the series of reactor vessels. Although the latter
embodiment permits somewhat longer catalyst life, the former
embodiment requires much smaller capacity heat exchangers. The process
considerably reduces catalyst usage as compared to a
conventional fixed bed epoxidn. process where all of
the catalyst is replaced or regenerated at the same time.
ACCESSION NUMBER: 130:66885
ITITLE: Epoxidation process using serially connected cascade
of fixed bed reactors
INVENTOR(S): Poxidation process using serially connected cascade
of fixed bed reactors
INVENTOR(S): Arco chemical Technology, L.P., USA
U.S., 9 pp.
COODENT TYPE: Patent ASSIGNEE(S): Arco chemical Technology, L.P., USA
U.S., 9 pp.
COODEN USXXAM
Patent
LANGUAGE: Patent
LANGUAGE: Patent
PATENT ASSIGNEE(S): Arco chemical Technology, L.P., USA
U.S., 9 pp.
COODENT TYPE: Patent
LANGUAGE: Patent
L

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

											LICAT						
											1997-						
C	2210	1427			21		1000	0701		CZ.	1998-	2210	427		î	0000	922
-	0022	1477			21		1000	0701		<u>س</u>	1998-	2310	46,		•	2200	023
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											, HR,						
											, LU,						
									SD,	SE	, SG,	SI,	SK,	SL,	ТJ,	TM,	TR
		TT,	UA,	υG,	υz,	VN,	YU,	ZW									
	RW:	GH,	GM,	KĖ,	LS,	MW,	SD,	SZ,	UG,	24	, AT,	BE,	CH,	CY,	DE,	DK,	ES.
		FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NI	., PT,	SE,	BF,	ВJ,	CF,	CG,	CI
		CM,	GA.	GN,	GW,	ML.	MR.	NE.	SN.	TI	, TG						
AL	9910	260			A		1999	0712		ΑU	1999- 1998-	1026	0		1	9980	923
E	1047	681			A1		2000	1102		EP	1998-	9526	31		1	9980	923
21	1047	681			B1		2003	1210									
		BE,															
RE	9813	1934		20,	n .	٠.,	2000	1121		RR	1998-	1383	4		1	9980	923
.76	2001	5262	80		7		2001	1218		7.0	1998- 2000- 2000-	5254	na .		ī	9980	923
	2205	101	••		ċ		2001	0527		D 11	2000-	1150	67		î	9999	923
	2000	101	006				2003	0127		~	2000-	DR 5.0	06		•	0000	616
RIORI1	Y APP	LN.	INFO	• •						US	1997-	9952	39		A 1	9971	219
										wn.	1998-	ED60	59	1	w 1	9980	923

THERE ARE 11 CITED REFERENCES AVAILABLE FOR

ANSWER 15 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN

A method for reacting an organic compound with a hydroperoxide
comprises ≥1 of steps 1-3: (1) reacting the hydroperoxide
with the organic compound to obtain a mixture composed of a reacted with the organic compound to occasion model organic compound organic compound and unreacted hydroperoxide; (2) separating the unreacted hydroperoxide from the mixture from step (1); (3) reacting the hydroperoxide separated in step (2) with the organic compound Thus, a tube reactor packed with strands of TS-1 was charged with 40 weights H2O2, MeOH, and propene at 40° and 20 bar to give 98.48 H2O2 conversion and 80.38 propene oxide selectivity.

ACCESSION NUMBER: 2000:117006 CAPLUS DOCUMENT NUMBER: 132:151672

Method for reacting an organic compound with a

Method for reacting an organic compound with a hydroperoxide: Harder, Wolfgang; Reach, Peter; Rieber, Norbert; Ruppel, Wilhelm; Teles, Joaquim Henrique; Walch, Andreas; Wenzel, Anne; Zehner, Peter BASF A.-G., Germany INVENTOR(S):

PATENT ASSIGNEE (S): PCT Int. Appl., 35 pp. CODEN: PIXXD2 Patent SOURCE:

DOCUMENT TYPE:

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

										APPLICATION NO.								
WO	2000007965				A1		20000217		WO 1999-EP5740						19990809			
	W:	AL,	ΑU,	ВG,	BR,	BY,	CA,	CN,	CZ,	GE	, HR	, HU,	ID,	IL,	IN,	JP,	KR,	
		ΚZ,	LT,	LV,	MK,	MX,	NO,	NZ,	PL,	RC	, RU	, SG,	SI,	SK,	TR,	UA,	US,	
		ZA.	AM.	AZ.	KG,	MD,	TJ.	TH										
	RW:	AT.	BE.	CH,	CY,	DE,	DK,	ES,	FI.	FF	, GB	. GR.	IE,	IT.	LU,	MC.	NL.	
			SE															
DE	1983				A1		2000	0217		DE	1998	-1983	5907		1	9980	807	
	2339																	
	9956																	
	1102															9990		
	1102				B1			0407									•••	
										GR	. IT	. I.T.	LU,	NI.	SE.	MC.	PT.	
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GT.	2002									.TD	2000	-5636	500		1	9990	909	
AT	2002 2637 2220	44	-		÷								119			9990		
FR	2220	106			· • 3								119			9990		
200	2001	DB 011	248		7			1031					48			0010		
	2001							0206					3			0010		
	2001							1006					, 13					
	6479		1/3										90					
								1112					:90 15907			9980		
PRIORITY	AFF.	ш	LINEO	• •							1000	- 1500		•	•	<b>3360</b>		

CASREACT 132:151672
9 THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

L7 ANSWER 16 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN (Continued) RECORD. ALL CITATIONS AVAILABLE IN THE RE

FORMAT

REFERENCE COUNT: THIS

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ANSWER 17 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
The catalytic activity of a titanium-containing heterogeneous catalyst
such as titania-on-silica which has been used to catalyze olefin epoxidn.
is effectively restored by washing the catalyst with water,
alc., ether, nitrile, ester, aromatic hydrocarbon, or ketone. Thus, a
non-silylated titania-on-silica catalyst was prepared in
accordance with the procedures described in U.S. patent number 3,923,843
  and
                            used to catalyze the epoxidn. of propylene with ethylbenzene hydroperoxide in a fixed bed reactor for 526 h. After removal from the reactor, the used catalyst gave only 231 conversion and 901 selectivity to epoxide in a batch epoxidn. of 1-octene with ethylbenzene hydroperoxide (1 h, 90°). A portion of the deactivated catalyst was washed with various solvents, dried in a 120° vacuum oven, and tested in batch epoxidn. of 1-octene using ethylbenzene hydroperoxide to give 62% conversion and 86% epoxide selectivity after washing with a-methylbenzyl alc. and 66% conversion and 86% epoxide selectivity after washing with deionized water.

SSION NUMBER: 1998:479458 CAPLUS
MENT NUMBER: 129:110418
E: Regeneration of heterogeneous catalyst for
                                                                                                                                    129:110418
Regeneration of heterogeneous catalyst for olefin epoxidation Carcoll, Kevin M.; Morales, Edrick; Han, Yuan-zhang Arco Chemical Technology, L.P., USA; Arco Chemie Technologie Nederland B.V. PCT Int. Appl., 37 pp. CODEN: PIXXD2 Patent English 1
    ACCESSION NUMBER:
  INVENTOR (S):
PATENT ASSIGNEE (S):
  SOURCE:
     DOCUMENT TYPE:
      LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                                                                                                                                                                                                                                  APPLICATION NO.
PATENT NO. KIND DATE APPLICATION NO. DATE

WO 9828072 Al 19980702 WO 1997-EP6625 19971127

W' AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, ID, IL, 19, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MK, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, AU, UG, UZ, VN, YU, ZW

RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SZ, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG

US 5916835 A 19990629 US 1996-770821 19961220

CA 2275264 Al 19980702 CA 1997-2275264 19971127

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EP 948403 Al 19991013 EP 1997-951993 19971127

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ER 9713751 A 20000201 BR 1997-13751 19971127

CN 1246075 A 20000301 CN 1997-181768 19971127

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PRIORITY APPLN. INFO: WO 1997-EP6625 W 19971127
                                PATENT NO.
                                                                                                                                                                         DATE
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                                                                                                                                                                            THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE
    REFERENCE COUNT:
      FORMAT
                             ANSWER 18 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN Propylene oxide (I) is manufactured by epoxidn. of propylene by using ethylbenzene hydroperoxide (II) ethylbenzene (III) solution, which is prepared by liquid phase autoxidn. of III. The
    give
an oil, (2) distilling of the oil to give condensed II solution in III to,
epoxidizing
of propylene (IV) by the II solution in the presence of Ti-containing
catalysts to give I associated with III, 1-phenylethanol, etc., (4)
removing IV by evaporation, and (5) evaporating the residual reaction
solution after
addition of 0.1-100% (based on I) C2-4 alcs., C2-8 ethers, C3-9 ketones,
C2-3
                                nitriles, and/or C6-7 hydrocarbons for separation of I-rich components
  nitriles, and/or C6-7 hydrocarbons for separation of I-rich components obtained
from the top and other components on the bottom. Thus, washing 3000 g II solution in III by aqueous NaOH, evapog. the solution in vacuo, epoxidizing IV by
the concentrated solution in the presence of Ti/SiO2 in fixed-bed reactor, removing IV, adding 0.05% PrOH, and evaporating gave I at selectivity 92% and loss of I $0.01%.

ACCESSION NUMBER: 198:108130 CAPLUS
DOCUMENT NUMBER: 128:167815
TITLE: Manufacture of propylene oxide with reduced loss of the product
INVENTOR(S): Yoshid, Masayuki; Oku, Noriaki; Ishino, Masaru
SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.
CODEN: JKXXAF
DOCUMENT TYPE: Japanese
FAMILY ACC. NUM. COUNT: 1
Japanese
FAMILY ACC. NUM. COUNT: 1
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APPLICATION NO.

JP 1996-210085 JP 1996-210085

DATE

19960808

19960808

DATE

19980217

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

JP 10045737 PRIORITY APPLN. INFO.:

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ANSWER 19 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN
A process is described for conducting an exothermic reaction between
22 reactants in a fixed-bed catalytic
reactor under isothermal conditions. The reaction mixture includes
21 compound having a lower b.p. than the other compds. present, the
quantity present being sufficient to consume, by vaporization, the heat
generated by the exothermic reaction. The reactor is operated
at the boiling pressure of the reaction mixture and facilitates a
rrent
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downflow of the liquid and the gas phases, the gas phase comprising the
vaporized component having the lowest b.p. Thus, the process was used
preparation of propylene oxide by catalytic oxidation of propene with ethylbenzene hydroperoxide using a Ti catalyst.

ACCESSION NUMBER: 1989:517684 CAPLUS
                                                                               1989:517684 CAPLUS
111:117684
                                                                               111:117684
Exothermic reaction process in a fixed-
bed catalytic reactor
Terlouw, Teunis; Van Os, George; Bakker, Anke Gezina
Shell Internationale Research Maatschappij B. V.,
 TITLE:
 INVENTOR(S):
PATENT ASSIGNEE(S):
                                                                                Neth.
Eur. Pat. Appl., 8 pp.
CODEN: EPXXDW
Patent
SOURCE:
 DOCUMENT TYPE:
LANGUAGE:
                                                                                English
  LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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                                                                                                                                     . APPLICATION NO.
                 PATENT NO.
                                                                                 KIND
EP 323663
EP 323663
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EP 323663
EP 323663
R: BE, DE, ES,
CA 1328273
ES 2030459
AU 8226901
AU 612741
CN 1033752
CN 1013552
JP 01203035
BR 8806626
ZA 8809400
PRIORITY APPLN. INFO::
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19900829
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BR 1988-6626
ZA 1988-9400
GB 1987-29555
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DATE

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19881125 19881213 19881215

19881215

19881215 19881215 19881215 A 19871218

L7 ANSWER 17 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN

ANSWER 20 OF 20 CAPLUS COPYRIGHT 2007 ACS on STN Alkenes; e.g. propene (I); were epoxidized in the liquid phase by means

Alkenes; e.g. propene (I); were epoxidized in the liquid phase by means an organic hydroperoxide, e.g., cumene hydroperoxide (III), ethylbenzene hydroperoxide (III), or tertbutyl hydroperoxide (IVI), introduced at several points of the reaction zone or in several fractions, and in the presence of Mo or V catalysts, using tert-buoN as solvent. This process may be carried out continuously in a tubular reactor or discontinuously in a vessel provided with a stirrer. Thus, 1500 g./hr. I was treated continuously at 90° with 480 g./hr. II, 480 g./hr. I was treated continuously at 90° with 480 g./hr. 1480 g./hr. tert-buoN and 16 g./hr. No naphthenate containing 3% Mo (V), in a stainless steel tube (1.5 m. long, internal diam 23-13 mm). Addhl. II was introduced at 2 points in the reaction zone at the rates of 520 and 529 g./hr. The hydroperoxide conversion was of 78.8% with a selectivity of 72.6% in propylene oxide (basis III), 56% II being recovered as cumyl alcohol. The following results were obtained by operating discontinuously 1 hr. at 90° with 0.16 part V (parts I, parts II, parts tert-BuoN, moles I-II consumed, % II converted, % epoxide selectivity, and % slochol recovery given): 5, 5, 16.9, 13.6, 94.0, 35.4, 97.0; 4.7, 4.7, 15.9, 6.4, 87.2, 81.6, 97.0; 7.0, 3.0, 15.1, 7.8, 86.2, 14.4, 81.0; 5.0, 10, 8.7, 2.1, 75.8, 62.3, 94. 1-Hexene (126 g.), 30 g. IV, 45 g. V, and 1.0 g. VI was heated at 110° in a discontinuous system, 5 g. IV added at 10°min. intervals up to an addnl. amount of 60 IV, and after 180 min. the mixture was distilled The hydroperoxide

IV, and after 180 min. the mixture was distilled The hydroperoxide conversion was 74% with a selectivity of 97% in epoxide (basis peroxide). Cyclohexene epoxide (VI) was also obtained in a discontinuous system by treating cyclohexene with III in the presence of V naphthenate, at 94 or 97% conversion with a selectivity in VII (basis III) of 96 or 87% after

or 60 min., resp.
ACCESSION NUMBER:
DOCUMENT NUMBER:
ORIGINAL REFERENCE NO.:

1966:412198 CAPLUS 65:12198 65:2222a-d Epoxidation of alkenes Halcon International, Inc. 13 pp. Patent Unavailable

ORIGINAL REFERENCE NO.:
TITLE:
PATENT ASSIGNEE(S):
SOURCE:
DOCUMENT TYPE:
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:

PATENT NO. APPLICATION NO. KIND DATE DATE NL 6507189 BE 665240 PRIORITY APPLN. INFO.: 19651216 NL 1965-7189 19650604 BE US 19640615

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